5 <u>PROCESS FOR PRODUCTION OF ESSENTIALLY</u> CHLORIDE-FREE CALCIUM SULFONATE

This patent application is a continuation-in-part of U.S. Patent Application 10/417,788 filed on April 17, 10 2003, the entire disclosure of which is hereby incorporated by reference.

FIELD OF THE INVENTION

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15 [0001] The invention relates to a process for the production of low base number essentially chloride-free calcium sulfonate.

BACKGROUND OF THE INVENTION

generally produced by the reaction of sulfonic acid with calcium hydroxide or calcium oxide, utilizing a promoter such as an alkanol. They can also be produced from sodium sulfonate by the use of calcium hydroxide or oxide and calcium chloride. Such sulfonates may be used as highly valued additives for lubricating oils such as passenger car, diesel, and marine engine lubricants. They may be further processed into overbased sulfonates, which have higher base numbers and are also used as additives for specialty lubricating oils.

[0003] When calcium sulfonate is derived from sulfonic acid, no chloride is needed, but the concentration of the final product is limited by the concentration of the sulfonic acid. In the case of natural petroleum sulfonic acid, concentration is typically less than commercially desired. Concentrating sulfonic acid itself is difficult due to its high corrosivity.

[0004] When calcium sulfonate is made from sodium sulfonate, chloride is required to make the reaction proceed. This leads to residual contaminating chloride in the final product. The sodium sulfonate is concentrated to the required concentration using a

solvent extraction process prior to conversion to the calcium product, since calcium sulfonate is more difficult to concentrate by this method.

[0005] A number of methods have been disclosed for the production of low base number calcium sulfonate.

[0006] US patent 5,804,094 teaches a method of producing a low base number calcium sulfonate of greater than 500 molecular weight using carboxylic acid and a high base number calcium sulfonate.

10 [0007] US patent 5,789,615 teaches the use of staged addition of calcium hydroxide to sulfonic acid to produce a low viscosity, low haze product without the use of promoters, especially without the use of chloride. The calcium hydroxide is added in two or more steps, with 30-15 180 minutes heat soak after each step.

[0008] US patent 4,615,841 describes a method of producing calcium sulfonates in the presence of an alkanol.

[0009] US patent 4,279,837 teaches the preparation of alkaline earth metal salts of alkyl benzene sulfonic acids by neutralization of the acid using an oxyalkylate as a promoter, thus also producing a chloride free calcium sulfonate.

[0010] US patent 3,719,596 describes a method of producing calcium sulfonate in which the reaction mixture is made acidic and then basic again using an alkanolamine.

[0011] US patent 2,779,784 teaches a method of producing calcium sulfonate in which sulfonic acid is neutralized with calcium hydroxide at 220 °F to 390 °F (104 °C to 199 °C), in the presence of 1/2 to 10 parts water per part calcium hydroxide. This would correspond to between 0.12 and 2.4 mol water per mol calcium hydroxide.

35 [0012] It would be advantageous to produce low base number calcium sulfonates, that are free of residual chlorine and easily concentrated, via a process suitable for use in a continuous reactor that can also produce

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products with a low viscosity.

SUMMARY OF THE INVENTION

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[0013] A method has been discovered to produce low base number calcium sulfonate, which is essentially free of residual chlorine and easily concentrated. The method can also produce a low viscosity product. The method may also be practiced in a continuous manner.

[0014] Accordingly the present invention provides a process for the production of low base number calcium sulfonates comprising:

- a. preparing a sulfonic acid-oil solution by adding about 1 to about 20 volumes of a miscible solvent to a sulfonic acid-oil feedstock and optionally removing dissolved or entrained SO₂ or SO₃ if present;
- b. mixing the sulfonic acid-oil solution with about 1 to about 5 moles of water per mol of sulfonic acid and about 1 to about 10 moles of calcium hydroxide per mole of sulfonic acid to provide a reaction mixture;
- c. heating the reaction mixture to a temperature in the range of about 40 °C to about 200 °C;
- d. separating excess calcium hydroxide from the heated-reaction mixture to produce a reaction product comprising solvent, oil, and calcium sulfonate;
- e. removing the solvent from the reaction product to produce an intermediate product comprising oil and calcium sulfonate;
- f. optionally concentrating the intermediate product by removing at least a portion of the oil to produce a concentrated product; and
- g. recovering the intermediate product and/or concentrated product, wherein the product is essentially chloride free calcium sulfonate in oil.

BRIEF DESCRIPTION OF THE DRAWINGS

[0015] Fig. 1 - This figure shows a flow chart of a continuous process for producing calcium sulfonate.

[0016] Fig. 2 - This figure shows the relationship

between the Strong Base Number (SBNC) of the calcium

sulfonate solution produced by the invention and the SBNC

of the product after solvent stripping.

[0017] Fig. 3 - This figure shows the relationship between product viscosity and the SBNC of the product after solvent stripping.

DETAILED DESCRIPTION OF THE INVENTION

The present invention provides a process for [0018] the production of low base number essentially chloridefree calcium sulfonate. In the context of the instant application, a low base number calcium sulfonate has a base number of 0 to about 50. By "essentially chloridefree" is meant a maximum chlorine content of 1000 ppm. In a particular embodiment, the present invention provides a process for the production of calcium sulfonate which comprises preparing sulfonic acid solution by adding about 1 to about 20 volumes of a miscible solvent to sulfonic acid and removing dissolved or entrained SO2 or SO₃ if present, mixing the resultant sulfonic acid solution with about 1 to about 5 moles of water per mol of sulfonic acid and about 1 to about 10 moles of calcium hydroxide per mole of sulfonic acid to prepare a reaction mixture, heating the reaction mixture to between about 40 °C and about 200 °C for a period of time up to about 60 minutes with stirring, separating excess calcium hydroxide and calcium salts of mineral acid from such a reaction mixture, and recovering solvent and oil to make an essentially chloride-free calcium sulfonate product. Sulfonic acid in an oil/solvent solution or [0019] dispersion is neutralized by calcium hydroxide in the

dispersion is neutralized by calcium hydroxide in the presence of a specific amount of water. Excess hydroxide and inorganic salt if any are subsequently removed from the reaction mixture by a suitable means such as centrifugation or filtration before removal of the

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solvent. In one embodiment, after removal of the solvent, the calcium sulfonate in oil may be concentrated by suitable means such as vacuum flashing or vacuum distillation, to produce a final product with a base number between 0 and about 50, and the desired final concentration.

The sulfonic acid in oil utilized may be [0020] The oil used in the process derived from petroleum oil. can be any suitably refined crude distillate. An example of a suitable feedstock is a vacuum distillate of appropriate molecular weight that has been refined by solvent extraction and/or hydrotreating to reduce the polynuclear aromatics content. The sulfonic acid solution used in the process is created by reacting the refined crude distillate with fuming sulfuric acid (about 27% - about 33% SO₃; oleum) or gaseous SO₃. When the feedstock is contacted with fuming sulfuric acid, monoaromatics are converted into mono-sulfonic acid and the residual poly-nuclear aromatics are converted into polysulfonic acid. The polysulfonic acid plus SO3 depleted sulfuric acid form a sludge. This reaction mixture is suitably diluted with about 1 to about 20 volumes of a miscible solvent to reduce viscosity, and the sludge is separated out by gravity settling, leaving the sulfonic acid in a solvent/oil solution. Dissolved or entrained SO₃ and/or SO₂, produced as a byproduct of side reactions between the oil and the SO_3 , are removed from the solution if present. One method of removal is stripping with nitrogen or another inert gas. The solution can also be centrifuged to remove traces of sludge prior to removal of dissolved or entrained SO2 or SO3.

[0021] Suitable solvents include any C_3 to C_{10} alkane, toluene or any low viscosity, miscible solvent. Most preferred is heptane or commercially available mixtures of heptane isomers.

[0022] To the cleaned sulfonic acid/solvent/oil solution is added about 1 mol per mol to about 5 mol per mol sulfonic acid of water and about 1 mol per mol to about 10 mol per mol sulfonic acid of calcium hydroxide

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to form the reaction mixture.

The reaction mixture is heated with mixing such [0023] as stirring to a temperature of from about 40 °C to about 200 °C, preferably from about 80 °C to about 120 °C. mixture is preferably stirred for a period of time up to about 60 minutes, more preferably up to about 30 minutes. The resulting mixture is then separated to remove excess calcium hydroxide and optionally if present, salts formed from residual sludge or SO2. 10 method of separating the mixture is centrifugation. Centrifugation should be performed for a sufficient amount of time to remove the excess calcium hydroxide and This period of time can be any such any salts. sufficient amount of time, for example, 20 minutes. presence of the solvent greatly improves the speed of 15 separation. The solvent is recovered from the clear centrate for recycle by any convenient means such as a solvent stripper. The product may be further concentrated via distillation or vacuum flashing to remove a portion or all of the unreacted oil. 20 recovered essentially chloride free calcium sulfonate product can be in various amounts of oil depending on the degree of concentration when the oil is removed. calcium sulfonate product in oil preferably has a viscosity of between 10 cSt/100°C and 100 cSt/100°C. 25 is preferable to remove the dissolved or entrained SO2 or SO3 if present from the sulfonic acid solution to obtain such viscosity. The essentially chloride free calcium sulfonate in oil can be recovered from the solvent removal step or after concentrating by removing at least 30 a portion of the oil or from both, for example, by collecting the essentially chloride free calcium sulfonate in oil at suitable steps in the process such as after the solvent removal step.

35 [0025] In one embodiment of the process of the invention, such process may be operated in a continuous fashion in a manner such as that shown in Fig.1.

Sulfonic acid 1 is added to a reactor 7, followed by

water 3 and lime 5. The resultant mixture then undergoes separation 9, with the lime and water being removed. The next step is solvent recovery 11, followed by concentration 13 to produce the calcium sulfonate in base oil 15.

[0026] The following examples are meant to further illustrate the invention without limiting its scope.

Comparative Examples - Set I

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[0027] A sulfonic acid solution (75 g) containing a mixture of petroleum sulfonic acid (8 wt%, average molecular weight of about 440 g/mol), commercial heptanes (60 wt%), and lubricating oil (32 wt%) was used in the following examples. This mixture was further treated by centrifugation and nitrogen stripping before being used in the examples.

[0028] Water, calcium hydroxide and tertiary butyl alcohol (TBA), as a promoter, were added to 75g of sulfonic acid. The resulting reaction mixture was heated with stirring for a specified time in an Erlenmeyer flask equipped with a reflux condenser. For temperatures above the boiling point of the mixture, a stainless steel reaction vessel was used to contain the mixture under pressure. After stirring, the mixture was transferred to a centrifuge tube and centrifuged for 10-20 minutes.

Table I shows the resulting Strong Base Number (SBNC, measured according to ASTM D974) of the centrate for various values of pretreatment, TBA content, water content, lime content, reaction time, reaction temperature, and centrifugation time.

- WITH PRETREATMENT AND WITH TBA COMPARATIVE EXAMPLES SET I TABLE I:

Centrifugal Centrate SBNC, time, min mg KOH/g	2.2	2.3	2.9	3.2	3.0	2.9
Centrifugal time, min	10	10	10	10	10	10
Reaction (time, min	10	30	30	30	30	30
Reaction Temp, *C	80	82	140	140	140	140
Lime, mol/mol sulfonic acid	4.0	4.0	4.0	4.0	4.0	4.0
Water, mol/mol sulfonic acid	2.4	2.4	1.7	1.9	1.7	2.7
TBA, mol/mol sulfonic acid	2.1	2.1	1.0	1.2	1.2	1.2
Comp. Example	ᠳ	2	m	4	ഹ	9

[0029] As can be seen, a base number of up to 3.2 can be obtained by optimizing the amount of TBA, water, and temperature.

Comparative Examples - Set II

- 5 [0030] The Comparative Examples in Set II were performed as in Comparative Examples Set I, however, the sulfonic acid was not treated by centrifugation and nitrogen stripping prior to reaction and no TBA was added. The results from these examples are in Table II.
- 10 Acidic results are shown as a negative SBNC value.

- NO PRETREATMENT, NO TBA COMPARATIVE EXAMPLES SET II TABLE II:

Centrate SBNC,mg KOH/g	-7.5	-0.3	-1.1	0.2
Reaction time, Centrifuge time, min	10	10	10	10
Reaction time, min	10	10	10	30
Reaction Temp, °C	7 7 8 9 8 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9	26	82	82
Lime, mol/mol sulfonic acid	3.0	3.0	3.0	4.0
Water, mol/mol Lime, mol/mol sulfonic acid sulfonic acid	0.5 4.6	12.7	0.5	2.7
Comp. Example	7 8	σ	10	11

[0031] These Comparative Examples show the results obtained without pretreating the sulfonic acid.

Comparative Examples Set III

- 5 [0032] The Comparative Examples in Set III were performed as in Comparative Examples Set I, however, the sulfonic acid was not treated by centrifugation and nitrogen stripping prior to reaction. The results from these examples are in Table IV. Acidic results are shown
- 10 as a negative SBNC value.

TABLE III: COMPARATIVE EXAMPLES SET III - NO PRETREATMENT, WITH TBA

Centrate SBNC, mg KOH/g	-7.0	1.0	8.0	0.8	-0.3	1.9	2.3	2.5	2.4
Reaction Reaction time, Centrifuge time, Centrate SBNC, Temp, °C min min min mg KOH/g	10	10	20	10	10	10	10	20	10
Reaction time, min	10	10	20	10	10	10	10	20	10
Reaction Temp, °C	26	26	26	26	26	26	09	09	82
Lime, mol/mol sulfonic acid	3.0	3.0	4.0	3.0	3.0	3.0	3.0	4.0	3.0
TBA, mol/mol Water, mol/mol sulfonic acid sulfonic acid	0.5	5.7	3.3	6.3	3.1	5.0	4.1	3.9	2.1
TBA, mol/mol sulfonic acid	10.9	1.4	3.3	7.5	7.9	4.5	4.6	4.2	2.1
Comp. Example :	12	13	14	15	16	17	18	19	20

[0033] These examples show results obtained without pretreating the sulfonic acid, but adding TBA to the reaction mixture. A maximum SBNC value of 2.5 was obtained.

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Examples

[0034] A sulfonic acid solution (75 g) containing a mixture of petroleum sulfonic acid (8 wt%, average molecular weight of about 440 g/mol), commercial heptanes (60 wt%), and lubricating oil (32 wt%) was used in the following examples. This mixture was further treated by centrifugation and nitrogen stripping before being used in the examples.

15 [0035] Water and calcium hydroxide were added to 75g of the treated sulfonic acid solution. The resulting reaction mixture was heated with stirring for the reaction time in an Erlenmeyer flask equipped with a reflux condenser. For temperatures above 82 °C, a 20 stainless steel reaction vessel was used to contain the mixture under pressure. After stirring, the mixture was transferred to a centrifuge tube and centrifuged for 10-20 minutes. Table IV shows the resulting Strong Base Number (SBNC, measured according to ASTM D974) of the centrate for various values of water content measured in 25 mol/mol of sulfonic acid, lime content measured in mol/mol of sulfonic acid, reaction temperature measured in °C, and reaction time and centrifugation time measured in minutes. Fig. 2 shows the correlation between the

SBNC of the centrate and the concentrated product.

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TABLE IV: EXAMPLES

Centrate SBNC, mg KOH/g		
Centrifuge time, min		10000 1000 10000
Reaction time, min	30000000000000000000000000000000000000	30 30 30 30 30 30
Reaction Temp, °C	60 60 60 60 82 117 117 117 117 117	1 1 1 1 1 4 0 0 0 0 0 0 0 0 0 0 0 0 0 0
Lime, mol/mol sulfonic acid	4 4 4 4 4 4 4 4 4 4 1 1 1 1 4 9	4 4 4 4 4 6 0 0 0 0 0 0
Water, mol/mol sulfonic acid		22.0 22.0 3.22 1.32
Example	1 2 2 4 3 9 7 8 9 7 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	11 11 11 11 10

[0036] These examples show that a base number of 3.6 can be achieved with the method of the invention. Table V and corresponding Fig. 2 show the relationship between the centrate SBNC and the stripped centrate (i.e., after solvent removal) SBNC and TBN such that a value for the stripped product can be extrapolated from Fig. 2.

TABLE V: RELATIONSHIP BETWEEN CENTRATE SBNC AND STRIPPED CENTRATE SBNC AND TBN

e Stripped Centrate TBN ASTM D2896	0.00 1.24 1.28 1.56 2.12 2.44 3.56 5.67 9.16
Stripped Centrate SBNC ASTM D974	-2.47 0.21 0.17 0.40 0.75 1.33 1.18 1.18 1.18 2.61 4.29 8.36
Centrate SBNC ASTM D974	-1.23 -0.18 -0.10 0.00 0.16 0.21 0.37 0.41 0.46 0.85 1.07 1.67
Example	100 8 4 3 5 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1

is advantageous to produce a higher base number product while still maintaining the product in the low base number product range. From Fig. 2, it can be seen that a base number of 3.6 correlates to a stripped centrate SBNC of about 9.1. Fig. 3 shows the relationship between the base number of the stripped product of the invention and the viscosity of the product. From Table VI and Fig. 3 it can be seen that a viscosity of about 15 cSt at 100°C correlates to a stripped centrate SBNC of about 9.1.

Table VI: Relationship between SBNC of Stripped Product and Viscosity of Stripped Product

Example	Stripped Centrate SBNC ASTM D974	Stripped Centrate Viscosity/100°C ASTM D445
1	-2.47	22
2	-1.17	250
3	0.00	3000
4	0.50	191
5	1.00	121
6	1.68	82.2
7	2.64	47.1
8	4.59	25.5
9	6.08	15.9
10	7.27	15.6
11	8.36	14.8

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